PANOGA: a web server for identification of SNP-targeted pathways from genome-wide association study data

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Associate Editor: John Hancock

ABSTRACT

Summary: Genome-wide association studies (GWAS) have revolutionized the search for the variants underlying human complex diseases. However, in a typical GWAS, only a minority of the single-nucleotide polymorphisms (SNPs) with the strongest evidence of association are explained. One possible reason of complex diseases is the alterations in the activity of several biological pathways. Here we present a web server called Pathway and Network-Oriented GWAS Analysis (PANOGA) to devise functionally important pathways through the identification of SNP-targeted genes within these pathways. The strength of our methodology stems from its multidimensional perspective, where we combine evidence from the following five resources: (i) genetic association information obtained through GWAS, (ii) SNP functional information, (iii) protein–protein interaction network, (iv) linkage disequilibrium and (v) biochemical pathways.

Availability: PANOGA web server is freely available at: http://panoga.sabanciuni.edu/. The source code is available to academic users ‘as is’ on request.

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Supplementary information: Supplementary data are available at Bioinformatics online.

Received on July 19, 2013; revised on December 16, 2013; accepted on December 18, 2013

1 INTRODUCTION

Genome-wide association studies (GWAS) aim to discover a subset of single-nucleotide polymorphisms (SNPs) that are associated with the onset and progression of complex disease phenotypes at a genome-wide scale. These studies have produced extraordinary successes in the past few years, with numerous new genetic variants identified with levels of significance that were once thought impossible to reach (Vischer et al., 2012). Standard GWAS approaches focus on single SNP analysis, and hence, it is difficult to incorporate weakly associated SNPs (Ball, 2011). For multifactorial diseases, this incorporation is especially important because no particular variant on a particular gene may have a strong effect, but the combination of the multiple variants with small effects on the polygenic disease explains the overall susceptibility to disease (Cirulli and Goldstein, 2010). In this regard, pathway-based approaches to GWAS, in which prior biological knowledge is integrated into association studies and sets of genes are analyzed simultaneously for association with a disease, have been proposed recently (Peng et al., 2010).

Because the analysis of single variants within isolated genes is not informative enough to explain the underlying disease mechanisms, another recent trend to further mine GWAS data is to integrate protein–protein interaction (PPI) networks (Schadt and Björkegren, 2012). Following a systems biology approach, these methods evaluate the pairwise or collective effects of genetic variants, where the functional consequence of each susceptibility allele may interfere with the interaction between pairs of proteins. A few studies available to date on the identification of the molecular interaction networks associated with the GWAS genes indicated that this approach can help to discover formerly unknown mechanisms underlying human diseases at the molecular level (Bebek et al., 2012).

To better understand the biological processes underlying complex diseases, we also considered the functional effect of a typed SNP in GWAS. Although the DNA polymorphisms that change protein function can have significant consequences, other types of SNPs, such as synonymous SNPs, do not have such serious effects in disease development mechanism. Functionally important SNPs, such as those that change amino acids, splicing sites; those that lead to gain or loss of stop codon; those that result in frame shift; and those that are found in regulatory region, are priority targets in disease studies and large-scale genotyping projects (Calabrese et al., 2009). Hence, we decided that SNP functional knowledge is valuable information to strengthen our pathway and network-oriented GWAS analysis (PANOGA) method.

To support the steps from SNP functionalization into PPI network analysis to functional enrichment in a single program, we developed a web server for post-GWAS analysis called PANOGA. With its multifactorial basis, PANOGA has a good potential to decipher the combination of biological processes underlying disease. PANOGA has been tested on several complex diseases, including rheumatoid arthritis (Bakır-Gungor and Sezerman, 2011), intracranial aneurysm (Bakır-Gungor and Sezerman, 2013), epilepsy (Bakır-Gungor et al., 2013) and Behçet’s disease (Bakır-Gungor et al., 2012), and proved to be...
useful. Throughout the time, the method has evolved with our efforts (Bakir-Gungor and Sezerman, 2012). This web server presents the latest version of PANOGA.

2 METHODS

For a given SNP that was found significant in a GWAS, first, the functional scores are fetched from The Functional Single Nucleotide Polymorphism (F-SNP) (Lee and Shatkay, 2008) and SPOT (Saccone et al., 2010) servers. F-SNP obtains the functional consequence of a SNP from multiple tools at splicing, transcriptional, translational and post-translational levels. SPOT score takes into account the effect of a SNP on the gene transcript (e.g. non-sense, frameshift and 5'- and 3'-UTR), the impact of an amino acid substitution on the protein product, evolutionary conserved regions and all possible linkage disequilibrium proxy SNPs with r2 over a predefined threshold in a specific HapMap sample (Saccone et al., 2010). Second, PANOGA combines the functional score and the GWAS P-value of a SNP and obtains Pw-values. Among all known SNP/gene transcript designations, the SNP is associated with the gene on which the SNP has the most important functional effect. Third, PANOGA transfers Pw-values of SNP to its associated gene.

During the network-oriented steps of PANOGA, genes with Pw-values are mapped into a human PPI network. Next, PANOGA finds connected sub-networks of the PPI network that has high total significance of genotypic P-values of the disease-predisposing SNPs with respect to the controls. To check for the biological relevance of the identified sub-networks, PANOGA performs functional enrichment. Finally, PANOGA integrates the functional enrichments of the generated sub-networks. The overview of PANOGA methodology is presented in Supplementary Figure S1. The Web site is implemented in HTML/PHP; operations are made with Java on the server side.

3 SERVER DESCRIPTION

PANOGA web server accepts GWAS results including reference SNP ID (rs#) and genotypic P-value of association for each tested SNP, as shown in Figure 1A. It is important to note that the PANOGA web server does not require individual genotypes, odds ratio, minor allele frequency or confidence intervals computed in a GWAS, which can have ethical considerations.

PANOOGA outputs SNP-targeted pathway and gene tables (Supplementary Tables S1–S4), and customized pathway maps, as shown in Figure 1B. SNP-targeted pathway tables of PANOGA present the number of associated SNPs from GWAS, the number of regulatory SNPs among the disease-predisposing SNPs, the number of SNP-targeted genes and the number of sub-networks that this pathway is found to be statistically significant.

The gene table of PANOGA includes the genes that are part of a functionally important pathway and at the same time identified in a sub-network. For each such gene, this table presents gene symbols, Entrez gene ID, number of times found in sub-network, number of associated pathways, list of associated pathways, number of typed SNPs in GWAS, functional information of the typed SNPs in GWAS, SNP regulatory potential and number of regulatory SNPs.

Customized Kyoto Encyclopedia of Genes and Genomes pathway maps of PANOGA, as shown in Figure 1C, enrich the utility of PANOGA results. These pathway maps help the users to visualize affected genes along different routes within the pathway map.

4 CONCLUSIONS

PANOGA is a web server with multiple functions that support pathway and network-oriented SNP-phenotype association analysis in a single system without requiring any additional manual processes. PANOGA represents a feasible solution for the identification of SNP-targeted pathways (candidate causal pathways) to bridge the gap between GWAS and biological mechanisms of complex diseases.

Funding: The work of B.B.G. has been supported by the Abdullah Gül University Support Foundation (AGUV).

Conflict of Interest: none declared.

REFERENCES


